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Sequence Analysis of the Cupin Gene Family in *Synechocystis* PCC6803

JIM M. DUNWELL

ABSTRACT

The recently described cupin superfamily of proteins includes the germin and germinlike proteins, of which the cereal oxalate oxidase is the best characterized. This superfamily also includes seed storage proteins, in addition to several microbial enzymes and proteins with unknown function. All these proteins are characterized by the conservation of two central motifs, usually containing two or three histidine residues presumed to be involved with metal binding in the catalytic active site. The present study on the coding regions of *Synechocystis* PCC6803 identifies a previously unknown group of 12 related cupins, each containing the characteristic two-motif signature. This group comprises 11 single-domain proteins, ranging in length from 104 to 289 residues, and includes two phosphomannose isomerases and two epimerases involved in cell wall synthesis, a member of the pirin group of nuclear proteins, a possible transcriptional regulator, and a close relative of a cytochrome c551 from *Rhodococcus*. Additionally, there is a duplicated, two-domain protein that has close similarity to an oxalate decarboxylase from the fungus *Collybia velutipes* and that is a putative progenitor of the storage proteins of land plants.

INTRODUCTION

The cupin superfamily (Dunwell, 1998) of functionally diverse proteins has been designated recently to include the germin and germinlike proteins from plants, their duplicated two-domain relatives, including the seed storage proteins (Bäumlein et al., 1995), and a wide range of other enzymes and binding proteins from microbes, plants, and animals (Dunwell and Gane, 1998). The name cupin (from the Latin *cupa*, a small barrel or cask) is derived from the tertiary β -barrel element, which comprises either the central core of these proteins (e.g., oxalate oxidase) or one of a number of discrete domains (e.g., araC transcription factors). Characteristically, the cupin element of these proteins has two histidine-containing motifs, which together with other conserved proline and glycine residues make up the structural framework and the putative metal-binding active site (Gane et al., 1998). The two conserved motifs are separated by a variable region, usually 15–20 residues in length.

The aim of the present study was to identify and categorize all cupin sequences within a single bacterial genome, namely, that of the unicellular cyanobacterium *Synechocystis* PCC6803 (Kaneko et al., 1996). This organism serves as the prokaryotic model for studying plantlike oxygenic photosynthesis, and the intention of the present study was to provide a basis from which the proliferation of related plant cupins could

be examined systematically. For example, it is estimated that the *Arabidopsis* genome contains at least 12 germinlike proteins (GLPs) (Dunwell, 1998), in addition to a large number of other related cupins.

METHODS AND RESULTS

A series of detailed database searches was conducted using the gapped BLAST (Altschul et al., 1997) and BLOCKS (Henikoff and Henikoff, 1991) programs to identify those *Synechocystis* protein sequences that contain the two conserved histidine-containing motifs described by Dunwell and Gane (1998). These two motifs are part of the β -strands designated, respectively, C/D and G/H within the two β -barrel elements of the bean storage protein phaseolin (Lawrence et al., 1994). A major theme in the present analysis, and the probable reason that this gene family had not been identified previously, is that the region between the two motifs is variable in length, with a minimum distance of 15 residues for many of the bacterial proteins, increasing to around 20 for the germins and GLPs and >20 for the storage proteins. The maximum of 64 residues is found in a barley globulin (gi|421978). This variable region, which can tolerate a range of insertions, is equivalent to the D/F loop of the β -barrel structure.

The main result achieved in the present analysis was identification of a total of 12 cupin sequences, of which 11 are single-domain proteins, with one example of a two-domain structure (Fig. 1). Each of these 12 has the characteristic cupin two-motif arrangement, with the consensus of motif 1 being PG(X)₅HXH(X)₁E(X)₇G and that of motif 2 being G(X)₅PXG(X)₂H(X)₃N.

The sequences can be subdivided into several classes on the basis of a range of criteria and an analysis of their nearest neighbors according to a BLASTP search (Table 1). In an assessment of potential function, and considering first the 11 single-domain proteins, two of the sequences (gi|1001180, gi|1652486) are thought to be phosphomannose isomerases (PMIs), one (gi|1653678) to be a dTDP-4-dehydrohannose 3,5-epimerase, and one (gi|1651977) a dTDP-6-deoxy-L-mannose-dehydrogenase. The latter two are very similar in sequence and should probably both be considered as epimerases.

It should be noted that residues 61–129 of the PMI sequence gi|1652486 are identical to the sequence encoded by nucleotides 3–208 (with a frameshift correction at position 104) in the upstream region of gi|287460, a sequence including the *Synechocystis* *groES* and *groEL* genes (Lehel et al., 1993). Presumably, this partial PMI coding region was accidentally ligated to the other coding regions during the cloning procedure and then was not identified because of the frameshift introduced by a sequencing error.

Another sequence from which a function can be deduced with some reliability is gi|1652717. This has as its closest neighbor the *Escherichia coli* sequence gi|1176281, a member of the recently designated group

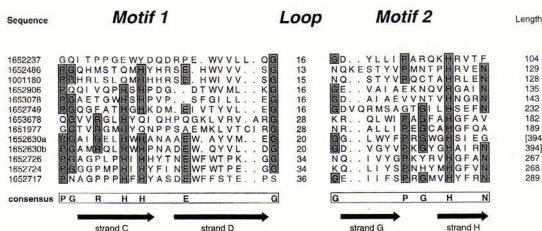


FIG. 1. Sequence alignment of the central core of cupin sequences from *Synechocystis* PCC6803. Sequences are denoted according to the GenBank gi identifier. The number followed by a and b denote the first and second domain in this two-domain protein.

SEQUENCE ANALYSIS OF CUPIN GENE FAMILY

TABLE 1. ANALYSIS OF CLOSEST NEIGHBORS FOR EACH OF THE CUPIN SEQUENCES FROM *SYNECHOCYSTIS* PCC6803^a

Sequence gi	Closest neighbor		Length AA	Identity %	Similarity %	Gaps %	Function
	gi	Species					
1652237	1657504	<i>E. coli</i>	47	34	55	—	Transcription regulator
1652486	1001180	<i>Synechocystis</i>	115	61	74	—	PMI
1001180	1652486	<i>Synechocystis</i>	102	63	78	—	PMI
1652906	1169942	<i>M. crystallinum</i>	95	27	50	9	GLP
1653078	347174	<i>Rhodococcus</i>	136	30	46	6	Cytochrome
1652749	1176281	<i>E. coli</i>	233	48	63	2	Pirin
1653678	141363	<i>S. enterica</i>	169	65	78	<1	Epimerase
1651977	1361427	<i>S. glaucescens</i>	179	42	55	—	Dehydrogenase
1652726	1652724	<i>Synechocystis</i>	233	48	68	—	?
1652724	1652726	<i>Synechocystis</i>	233	45	65	—	?
1652717	1652724	<i>Synechocystis</i>	132	26	47	6	?
1652630	1604990	<i>C. velutipes</i>	325	38	55	3	Oxalate decarboxylase

^aEstimated by use of the gapped BLASTP program, showing the length of the region of greatest similarity, the percentages of identical and similar residues, and the percentages of gaps needed to give maximum similarity (TBLASTN was used for the two-domain GI1652630 sequence, which has a DNA sequence as its neighbor).

of nuclear proteins, the pirins (Fig. 2) (Wendler et al., 1997). Additionally, gi1653078 is very similar to the *Rhodococcus* sequence gi347174, a putative C551 cytochrome, and gi1657504 is similar to the *E. coli* transcriptional regulator gi1657504. A lower degree of similarity, with gaps, is found between gi1652906 and its nearest neighbor, the GLP (gi1169942) from *Mesembryanthemum crystallinum*.

Although no function can be assigned yet to the remaining single-domain sequences, the two-domain 394-AA protein (gi1652630) may be an oxalate decarboxylase, as predicted from its similarity (Fig. 3) to the 447-AA oxalate-degrading enzyme (Mehta and Datta, 1991) from the wood-rotting fungus *Collybia velutipes*. Its sequence (gi1604990) has been published recently (Datta et al., 1996).

A detailed quantitative analysis of the cupin sequences shows a number of interesting features. If the single-domain proteins are considered first, in general terms the distance between motifs increases in line with the in-

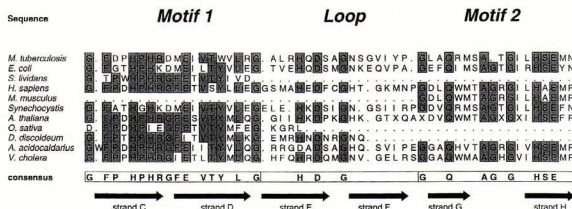


FIG. 2. Sequence alignment of the central core of pirins from a range of microbes, plants, and animals. The details of the sequences used are as follows: *Mycobacterium tuberculosis* (gi2213518), *Escherichia coli* (gi1789847), *Streptomyces lividans* (translation of part of the -ve strand of the actinophage phi C31 attachment site gi48953), *Homo sapiens* (gi1907076), *Mus musculus* (EST gi1282795), *Synechocystis* (gi1652749), *Arabidopsis thaliana* (EST gi950773), *Oryza sativa* (EST gi1631547), *Dityostelium discoideum* (-ve strand 302-3 of upstream sequence of *spiA* gene, gi1177288), *Alicyclobacillus acidocaldarius* (manually edited from -ve strand upstream sequence of amylase gene gi39300), *Vibrio cholerae* (sequence included in 538-nucleotide unfinished fragment gnlTIGR/GVCCX37R).

		Motif 1	
I25120	111	FSFSKQRL--QTGGWARQQNEVVLPLATNLACTNMRLEA GAIRELHWHKN-AEWAYVLKG	167
		++FSK L GG +Q P++ +A M LE GAIRELHWH N AEWAYV++G	
Synec.	56	YAFSKTPLVLYDGGTTKQVGTYNFPGSKMGAGVYMSLE GAIRELHWHANA-AEWAYVMEG	115
		Motif 2	
I25120	168	STQISAVDNEGRNYISTVGF GDLWYFPFGIPHS LQATADDPGSEFILVDFSGAFNDDGT	227
		T+I+ EG+ I+ V G LWYFP G HS++ P+ ++F+LVF+ G F++ T	
Synec.	116	RTRITLTSPEGKVEIADVDK GGLWYFPRGWGHS IEGIG--PDTAKFLLVFNDGTFSEGAT	173
I25120	228	FLLTDLWSHVPMEVALKNFRAKNPAWSHIPAQQLYIFPSEPPADNPDPVSPQG---TV	284
		F +TDWLSH P+ +N A + +P +Q+YI S PA +PGG +	
Synec.	174	FSVTDWLSHTPIAWVEENL-GWTAQAQVQLPKKQVYI--SSYGPASGPLASATPQGQTAKI	231
		Motif 1..	
I25120	285	PLPYSFNFSSVEPTQYSGGT-AKIADSTTFNISVAIAVAEVTVE GAIRELHWHPTDEW	343
		+P++ N +P GG ++A+ F S + A + +EPQA+R+LHWH DEW	
Synec.	232	EVPHHTNLLGQQPLVSLGGNELRLASAKEFGSFMNTGALIHLE GAMROLHWHHPNADEW	291
		Motif 2	
I25120	344	TFEILGNAVRTIFAAQSVASTFDYQ GDIAYVPASMGHYVENI GNNTLTLYLEVENTDRFA	403
		+ + G +T+FA++ AS Q GD+ YVP GH + N L + VFN +	
Synec.	292	QYVLDEMDLTVFASEGKASVSR QGDVGVYVFKGYGHAI RNSQKPLDIVVFNDDGYQ	351
I25120	404	DVSLSQWLALTPPSVVQAHLNLDDETLEAEL	433
		+ LS WLA P SV+ + E +L	
Synec.	352	SIDLSTWLASNPFSSVLGNTFQISPELTKKL	381

FIG. 3. Protein sequences of the two-domain proteins from *Synechocystis* PCC6803 (gi|1652630) and *Collybia velutipes* (I25120), compared by gapped BLASTP. Score = 210 bits (558), expect = 1e-53, identities = 123/330 (37%), positives (+) = 179/330 (53%), gaps = 11/330 (3%). The shaded boxes denote motifs 1 and 2 within each domain.

crease in overall size of the protein (Fig. 1), from a minimum of 13 residues in the 129-AA gi|1652486 to 36 residues in the 289-AA gi|1652717. The one notable exception to this trend is the 232-AA gi|1652749, which has a 16-residue intermotif loop. As reported, the two-domain sequence is related to a fungal decarboxylase. Both have 20 residues between motifs, a spacing characteristic of many of the GLPs from higher plants.

DISCUSSION

Before the present study, only four of the *Synechocystis* sequences had been identified as cupins (Dunwell and Gane, 1998). Three of these were single-domain proteins, namely, the PMI gi|1001180 and its two related sequences, gi|1652906 and gi|1653708. The other was the two-domain protein gi|1652630. The total number in this paralogous family is now increased to 12 (Fig. 1 and Table 1), although not all of these proteins fulfil the arbitrary definition used in the recent analysis of the *E. coli* genome (Blattner et al., 1997). These authors used the term to include all those ORFs that share at least 30% sequence identity over more than 60% of their lengths. However, it is probably not appropriate to apply this strict criterion to the present group of 12 sequences, which vary in length from 104 to 394 AAs and in which the two most conserved motifs are separated by a nonconserved region of variable length.

Four of the proteins identified in the present study have a possible functional connection, in that they are concerned with cell wall synthesis. The best known are the PMIs (EC 5.3.1.8), enzymes that catalyze the interconversion of mannose-6-phosphate and fructose-6-phosphate. Although they are considered to be zinc-containing metalloproteins, an Fe(III)-hydroxyphenylalanine site has been identified recently in the PMI from *Candida albicans* when expressed in *E. coli* (Proudfoot et al., 1996; Smith et al., 1997). On the basis of sequence comparison, PMIs have been divided into three classes (Proudfoot et al., 1994), within which

the class II enzymes (those described here) are involved in a variety of pathways, including capsular polysaccharide biosynthesis and D-mannose metabolism. Interestingly, in *Synechocystis* there seems to be no example of the bifunctional GDP-mannose pyrophosphorylase/PMI enzyme (the PMI domain of about 130 AAs is located at the C-terminus of the protein) found in some Archaea (e.g., the 448-AA gi|2649495 from *Archaeoglobus fulgidus*) and many eubacteria (e.g., the 428-AA gi|230580 from *Vibrio cholerae* and the 470-AA gi|2313118 from *Helicobacter pylori*) and thought to be involved, for example, in the polymerization of alginate, a viscous mucoid exopolysaccharide. Instead, the two functions in *Synechocystis*, as in *Methanococcus jannaschii* (J.M. Dunwell, unpublished observations), are carried out by two separate enzymes, the phosphorylase function by the 367-AA gi|1653346 and the isomerase function by the two PMIs (approximately 128 AA), gi|1001180 and gi|1652486, identified previously. Other bacteria, such as *E. coli*, contain a family of related enzymes, including PMIs (e.g., the 152-AA gi|147164), as well as several bifunctional enzymes (e.g., the 471-AA gi|585853, the 478-AA gi|1155018, and the 483-AA gi|1584629).

Enzymes, such as the dTDP-4-dehydrothiamine 3,5-epimerase (gi|1653678) identified here, are involved in the synthesis of bacterial exopolysaccharides. These enzymes include the ExpA8 protein recently shown to be involved in the synthesis of galactoglucan (exopolysaccharide II) in *Rhizobium melioli* (Becker et al., 1997) and the TDP-deoxyglucose epimerase (L33181) that is part of the biosynthetic pathway for ascaryle, a lipopolysaccharide component in *Yersinia pseudotuberculosis* (Thorson et al., 1994). The related dTDP-6-deoxy-L-mannose-dehydrogenase (gi|1651977), identified in this study as a cupin, is also involved in cell wall synthesis.

In contrast to the presumed function in cell wall synthesis assigned to the previous four sequences considered, the identity of sequence gi|1652749 seems to be as the nuclear protein pirin, one of a highly conserved group of proteins (Fig. 2) thought to interact with the nuclear factor I/CCAAT box transcription factor (NFI/CTF1) (Wendler et al., 1997). Of the other sequences, gi|1652237 has the transcriptional regulator gi|1657504 from *E. coli* as its closest neighbor, and gi|1653078 is similar to gi|347174, a cytochrome C551 gene from *Rhodococcus*.

Unfortunately, no function can be assigned to any of the other single-domain cupins listed in Figure 1, although it is hoped that as this superfamily is analyzed in more detail, the subgroups will be identified systematically by their homology to proteins of known function—a process that enabled the oxalate oxidase identity of the wheat gf 2.8 germin to be established (Lane et al., 1993).

The detailed analysis summarized in Figure 1 confirmed that all the cupin sequences found in *Synechocystis* contain the characteristic two-motif structure, with the intermotif distance varying in length from 13 (a uniquely short spacing for this superfamily) to 34 residues. Despite the range of spacing found in this study, it is interesting to note that none of the single-domain cupins in *Synechocystis* has an intermotif distance of 20 residues, the spacing found in the two-domain sequence gi|1652630. There are a number of possible explanations for this peculiarity. First, it may be that the duplication occurred in a protein with a 16-residue spacing, followed by the insertion of 4 residues in each domain; this seems unlikely. Alternatively, the 20-residue progenitor may have been lost through natural selection or may simply not have been identified in this study. Again it seems unlikely that any other cupin sequences in this genome remain undiscovered. In a further attempt to find close relatives to the two-domain 20-residue protein from other organisms, the 57-AA sequence spanning the two motifs of domain 1 was used as a probe in a BLASTP search. This revealed an *Arabidopsis* GLP (U75207) as the closest neighbor, with the closest nonplant sequence being the hypothetical protein gi|2128971 from the archaeon *M. jannaschii* (37% identical, 48% similar over a distance of 51 AAs). However, this 125-AA protein has only a 16-residue gap. Similarly, use of an equivalent sequence from domain 2 revealed the 113-AA sequence gi|1881251 from *Bacillus subtilis* as the nearest neighbor (37% identical, 62% similar over 51 residues). This sequence also has a 16-AA gap. Therefore, to date, there is no known example of a single-domain, 20-AA gap protein from a prokaryote. This apparent lack of any progenitor 20 protein, allied to the multitude of prokaryotic two-domain proteins with the 20 spacing, suggests that there was only one 20 protocupin, which underwent a duplication to produce gi|1652630 and its equivalents in other species and left no extant progenitor (or at least none identified to date).

Comparison of the alignment of the two-domain sequences from *Synechocystis* and *C. velutipes* (Fig. 3) confirms that they are probably derived from the same progenitor and that the former sequence may be the direct progenitor of the latter. This view is reinforced by analysis not only of the conserved motif regions but also of the intervening intermotif regions. Specifically, there are several identical residues in the two

20-AA intermotif regions of the first domain of each protein (consensus XTXIXXXXXXEGXXXIXVXX) and a different set of identical residues (consensus XXXTXFAXXXXASXXXQX) in the intermotif regions of the second domain. This suggests strongly that there was divergence of the two domains of a precursor protein after duplication of a 20-residue protocupin, followed by further minor divergence during the postduplication phase, leading eventually to the present day sequences.

In terms of both their number and their range of size and the presence of a two-domain sequence, the evidence presented suggests that the spectrum of cupins found in *Synechocystis* is closer to that of higher plants rather than to that found in more primitive bacteria (to date, only two cupins have been identified in *M. jannaschii*) (Dunwell, 1997). Presumably, there was a rapid expansion of cupin diversity during the evolution of *Synechocystis*. This conclusion may be related to the observation that the genome of this organism contains 99 ORFs with similarity to transposases (Kaneko et al., 1996). It was suggested that this high frequency could be linked to frequent rearrangement of the genome during and after establishment of this species. More recently, Cassier-Chauvat et al. (1997) characterized three specific insertion sequences from *Synechocystis* and suggested, on the basis of homology, that they were spread through horizontal transfer between evolutionarily distinct organisms.

Identification of the prokaryotic two-domain sequence in the present study also complicates the conclusion reached by Bäumlein et al. (1995) that "the extant spherulins and germins might represent a stage of seed globulin gene evolution *before* [my emphasis] the domain duplication event had occurred."

If the linear structure of these 12 *Synechocystis* cupin sequences is considered, it can be seen that there is an overall increase in spacing between the motifs, coincident with the increase in length of the proteins (Fig. 1). If it is assumed that the progenitor of these proteins was the smallest and most simple of the protocupins, as the overall sequence grew in length by addition of residues at each end, this must have been accompanied by insertion of residues into the variable region, namely, the E/F loop at the end of the β -barrel (Gane et al., 1997). This gradual increase in complexity led eventually to addition of α -helical regions at each end of the protein, duplication into two long $\alpha + \beta$ domains, and finally to assembly of the protein subunits to give the trimeric quaternary structure found in the abundant storage proteins of land plants (Lawrence et al., 1994) (Table 2). In addition, by the present stage of the evolutionary process, two of the three conserved histidines have been lost in most storage proteins (all three in some examples), and as these residues are implicated as being catalytically active (Gane et al., 1998), it is likely that these multimeric proteins no longer have any enzyme activity (none is known). In this regard, it is also not known if the *Synechocystis* two-domain protein or any of the more primitive two-domain storage proteins show oxalate decarboxylase activity (cf. I25120).

Duplication of domains, followed by their subsequent divergence during evolution, is known to occur in other proteins, such as the zinc metalloenzyme glyoxylase 1, which catalyzes the glutathione-dependent inactivation of toxic methylglyoxyl (Cameron et al., 1997). In this example, sequence alignment showed that 13 of 74 residues were identical in domain 1 and 13 of 59 in domain 2. Such relatively low levels of identity occur presumably because of the small number of residues required to provide the conserved structural elements in such proteins. Undoubtedly, as analysis techniques become more sophisticated, more cases of

TABLE 2. CHARACTERISTICS OF A SAMPLE OF CUPIN PROTEINS FROM THE EVOLUTIONARY SEQUENCE FROM BACTERIA TO HIGHER PLANTS^a

Name	Species	Length AA	Histidines	Loop AA	Domains	Subunits
PMI	<i>Synechocystis</i>	128	3	13	1	1
GLP	<i>A. thaliana</i>	200	3	20	1	1
Oxalate oxidase	<i>T. aestivum</i>	201	3	23	1	75
Oxalate decarboxylate	<i>C. velutipes</i>	447	3	20	2	1
Vicilin	<i>P. vulgaris</i>	397	1	27	2	3

^aThe proteins are classified according to their name, species of origin, total length, number of conserved histidine residues in the two motifs, length of the intermotif loop, number of domains, and number of subunits in the mature protein.

this type of evolutionary process will be revealed, and the number of basic underlying 3D structures will be found to be restricted (Godzik, 1997).

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